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Synergism between non-thermal plasma and photocatalysis: Implications in the post discharge of ozone at a pilot scale in a catalytic fixed-bed reactor



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ABSTRACT

This study focuses on Butyraldehyde (C_4H_8O) degradation using catalytic ozonation oriented to the optimization of non-thermal plasma/photocatalytic coupled processes. After the first oxidation by non-thermal plasma coupled photocatalytic processes, a second oxidative step leading to the decomposition of the residual Butyraldehyde by Ozone is catalyzed by Mordenite, ZSM-5 and γ -alumina. The Butyraldehyde degradation in the presence of Ozone under operational parameters such as plasma energy, Butyraldehyde inlet concentration, relative humidity and the height of the catalytic bed are reported in detail. The optimum operating conditions found were: $Q_{air} = 1 \text{ m}^3 \text{ h}^{-1}$, $[C_4H_8O] = 100 \text{ mg m}^{-3}$, $SIE_{plasma} = 18 \text{ JL}^{-1}$, for Butyraldehyde degradation mediated by ZSM-5. The formation of intermediate by-products was monitored during the degradation of Butyraldehyde in the presence of Ozone and ZSM-5 catalyst to suggest a degradation pathway. The coupling of catalytic ozonation by non-thermal plasma/photocatalysis lead to the total decomposition of Ozone when catalyzed by ZSM-5and presents a positive effect on the mineralization capacity. The pollutants degradation intermediates were identified by GC-MS and the oxidative states on the catalytic bed were characterized by XPS.

1. Introduction

Outdoor air pollutants have common sources: industry, transportation, heating and agriculture [1], they affect negatively health and climate. Recently, European directive 2016/2284/EC has outlined the reduction of emissions by 2030 for air pollutants like NH₃, SO₂, NO_x, volatile organic compounds (VOCs) (excluding CH₄)) [2]. France, for example, is committed to reduce from 52% up to 77% its emissions of VOC, NO_x, SO₂, and by 13% ammonia by 2030 [3]. All the processes using a liquid phase combined with an oxi-reduction mechanism (Stretford, Ferrifloc, Sulfurex, Burner-Scrubber, Catalyst-Scrubber, Ozone Process, etc.) expensive equipment subject to corrosion [4]. This leads to high investment and maintenance costs, as well as large consumption of chemical reagents and environmental problems created by discharge of the waste effluent after treatment [5]. Processes not requiring corrosive reagents but a power supply and producing only

inorganic by-products (CO_2 , H_2O) are needed at the present time. Technologies "zero waste & zero reagent" coupling non-thermal plasma (NTP) and Dielectric Barrier Discharge (DBD) with catalytic oxidation look promising to achieve this objective at low energy consumption [6–9]. VOCs removal has been reported including isovaleraldehyde, isovaleric acid, trimethylamine, benzene, ammonia and dimethyl disulfide [10–15]. The amount of ozone produced during VOCs treatment via oxidation processes was observed to be high > 20 ppm [16]. Zeolites have been reported to induce a higher ozone decomposition compared with activated carbons, which are flammable under high concentration of ozone [17,18]. Thus, the goal of this study is to improve the removal of volatile organic compounds by coupling of DBD-plasma-photocatalysis and catalytic ozonation and its consequences on Butyraldehyde degradation.

The combination of DBD-plasma/photocatalysis/catalytic ozonation was determined to optimize the operating process for better removal of

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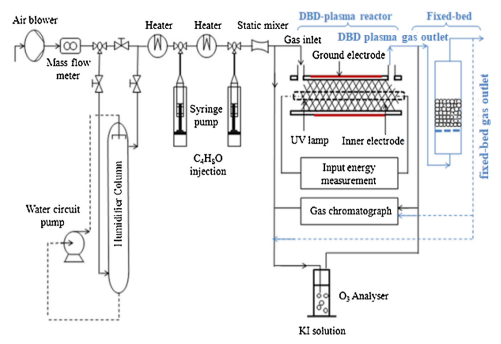


Fig. 1. Schematic of the experimental setup: annular and fixed-bed reactors.

malodors and air pollution. In this work, by adding the zeolite fixed bed reactor to the plasma/photocatalysis reactor we aim to: (i) increase the degradation of the residual butyraldehyde in the fixed bed, and (ii) reduce the amount of ozone created in the plasma/photocatalysis reactor.

2. Experiment and measurement

2.1. Experimental setup

The experimental setup used to treat VOCs under DBD-plasma/fixed-bed system is shown in Fig. 1. The setup includes a control for air stream generation system, two continuous pluck-flow reactors for oxidation step (annular and fixed-bed reactors) gas sampling equipment. The DBD-plasma reactor works in photocatalytic mode and/or in plasma mode or in simultaneous mode. The DBD-plasma reactor consists of two concentric cylinders, one outer cylinder of 76 mm and an inner cylinder of 58 mm with wall thickness of 4 mm. Both cylinder are 100 cm long and contain UV lamp (Philips TL 40 W/05). In the active zone of the reactor, between the inner wall of the outer cylinder/outer wall, polluted air stream is treated under UV-light (Ahlstrom Research and Services). The physical properties of the catalyst are given in Table 1.

Thus, UV lamp was placed in the inner concentric cylinder in order to have a homogeneous irradiation. To generate DBD-plasma, high voltage power is applied to the reactor which is covered by a copper grid forming the outer electrode and the inner was made out of Al. The applied voltage is delivered by a generator as a sinusoidal waveform up to 10 V and then amplified to 30 kV [9–11,16]. A detailed diagram of the DBD-plasma reactor was presented in Fig. 2, indicating the different experimental parts used to create the plasma. The air stream at the outlet of the DBD-plasma reactor, under goes a second oxidation step. The fixed-bed reactor consists of a Pyrex glass of 30 mm as internal

Table 1Main characteristics of TiO₂-GFT.

Туре	BET surface (m ² /g)	Crystal structure	Diameter (nm)
PC500	300	Anatase 100%	5-10

diameter and 15 cm length. At the bottom of the tube, sintered glass is placed to hold the catalyst, and the catalyst weight is varied between 20 and 60 g and the height of the catalytic bed varied from 5 to 15 cm (see Fig. 1).

2.2. Pollutants and Zeolites

Butyraldehyde (C_4H_8O , 99.9% purity) was purchased from Sigma Aldrich. It is injected continuously in liquid state by means of a programmable syringe pump system (KdScientific Model 100). A static mixer upstream ensures a good mixture of the gaseous effluent. Three catalytic materials were tested, namely two types of commercial zeolites from TOSOH Corp Tokyo75 Japan with microporous structures (ZSM-5 and Mordenite) and γ -alumina with mesoporous structures from Alfa Aesar Company. The characteristics of the materials are presented in Table 2.

2.3. Analytical device

The concentration of Butyraldehyde (BUTY) at the inlet/outlet of annular and fixed-bed reactor was sampled manually with a 500 µL syringe and analyzed by a gas chromatography (Fisons GC9000) equipped with a flame ionization detector (GC-FID). The various intermediate by-products produced during the Butyraldehyde degradation were measured in a Gas Chromatograph-Mass Spectrometer (GC-MS) coupled to an infrared (IR) detector. The Ozone concentration at the outlet of annular and fixed-bed reactor was monitored by iodometric titration method. The outlet air stream (annular reactor and/or fixed-bed) is bubbled through a potassium iodide (KI) solution with 10⁻² M concentration until a yellow color emerges. The sample is then titrated with sodium thiosulfate solution (Na₂S₂O₃) with 10⁻² M until a colorless solution is obtained [10]. The relative humidity in the air stream was controlled. All experiments were carried out at ambient conditions, atmospheric pressure and room temperature. The temperature and relative humidity (RH) were determined using a specific sensor (Testo 445). An AXIS NOVA photoelectron spectrometer (Kratos Analytical, Manchester, UK) provided for with monochromatic AlKa $(h\nu = 1486.6 \text{ eV})$ anode. Binding energies (BE) were calibrated against the standard C1s binding energy at 284.6 eV [19,20]. A Multipak (version 9 software) using 70:30 Gaussian: Lorentzian peak shape and a

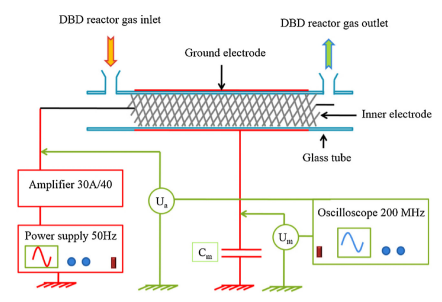


Fig. 2. General electric diagram of the DBD-plasma reactor.

Table 2 Main characteristics of the zeolites and γ -alumina.

Type	ZSM-5	Mordenite	γ-alumina
Pore diameter (Å)	(5,7 × 5,3)+	(6,6 × 7)+	150
	$(5,1 \times 5.5)$	$(3,5 \times 6,7)$	
Si/Al (mol/mol)	1880	220	0
BET surface (m ² /g)	308	509	237
Pore volume (cm ³ /g)	0,11	0,21	0,15
Binder type (% weight)	Alumina (20%)	Alumina (20%)	-
Pellet diameter (mm)	1,5	1,5	3

Shirley background function were used for Spectra deconvolution [21]. Process performance was evaluated by measuring: Inlet $[C_4H_8O_{inlet}]$ and outlet $[C_4H_8O_{outlet}]$ concentrations of C_4H_8O

Removal efficiency of C4H8O: RE (%)

$$= \frac{([C_4 H_8 O]_{inlet} - [C_4 H_8 O]_{outlet}) \times 100}{[C_4 H_8 O_{outlet}]}$$
(1)

Specific input energy: SIE
$$(J L^{-1}) = \frac{3600 * P(W)}{Q(m^3 h^{-1}) \times 1000}$$
 (2)

where P is input power adjusted by changing the applied voltage $(U_{\rm a})$ and Q is the flowrate.

$$Selectivity of CO: SCO (\%) = \frac{[CO]_{outlet} - [CO]_{inlet}}{4 \times RE (\%) \times [C_4 H_8 O]_{in}} * 10000$$
(3)

Selectivity of CO2:
$$SCO_2(\%) = \frac{[CO_2]_{outlet} - [CO_2]_{inlet}}{4 \times RE(\%) \times [C_4H_8O]_{inlet}} * 10000$$
 (4)

3. Results and discussion

3.1. Butyraldehyde and Ozone removal: Mordenite & $\gamma\text{-alumina supports Vs}\ ZSM\text{-}5$

Fig. 3 shows the concentrations of Butyraldehyde and Ozone within DBD-plasma/fixed-bed reactor in processes mediated by ZSM-5, Mordenite and γ -alumina. Regarding Butyraldehyde degradation, it was observed that the outlet concentration of Butyraldehyde decreases from 60.75 mg m $^{-3}$ (DBD-reactor gas outlet) to 41.23 mg m $^{-3}$ (fixed-bed gas outlet), 51.92 mg m $^{-3}$ (fixed-bed gas outlet) and 59.43 mg m $^{-3}$ (fixed-bed gas outlet) when using ZSM-5, Mordenite and γ -alumina, respectively, for C_4H_8O inlet concentration of 100 mg m $^{-3}$. A similar behavior

was observed for ozone. The Ozone concentration decreases from $104.79~mg~m^{-3}$ (DBD-reactor gas outlet) to $84.51~mg~m^{-3}$ (ZSM-5), $88.73~mg~m^{-3}$ (Mordenite) and $94.57~mg~m^{-3}$ (γ -alumina) with SIE of 8 JL $^{-1}$. When using γ -alumina, the presence of water vapor (HR = 45%) inhibits the decomposition of ozone and therefore negatively influenced Butyraldehyde oxidation. A competition between water, Butyraldehyde and Ozone on the sites of adsorption of γ -alumina seems to be responsible for this observation [22]. Only 11% of the Butyraldehyde and 14% of Ozone was removed in the fixed-bed reactor. Roscoe et al. [23], by FTIR (Fourier Transform InfraRed) spectroscopy reported the high affinity of γ -alumina for water molecules at sites that may be involved in the decomposition of ozone [23]. As it is readily seen in Fig. 3, ZSM-5 leads to highest Butyraldehyde degradation and Ozone oxidation.

3.2. Single Plasma and Catalytic Ozonation: case of ZSM-5

The effects of three major operating parameters, namely: (i) plasma specific energy, (ii) initial Butyraldehyde concentration and (iii) relative humidity on Butyraldehyde degradation using ZSM-5 were monitored. The gas flow rate $(1 \, \text{m}^3 \, \text{h}^{-1})$ and the weight of the catalyst (20 g) have been fixed as constant for all the experiments.

3.2.1. Effect of specific input energy

Two experiments with specific input energy of 9 J $\rm L^{-1}$ and 18 J $\rm L^{-1}$ have been carried out under relative humidity of 5 % and 100 mg m $^{-3}$ of Butyraldehyde. The evolutions of Butyraldehyde concentration at the outlet of the first reactor (DBD- plasma reactor gas outlet) and the second reactor (fixed-bed gas outlet) for both SIE are reported in Fig. 4. The Butyraldehyde oxidation is carried out for 8 hours. $\rm C_4H_8O$ oxidation was observed in DBD-plasma process and then moves through the catalytic bed for a second degradation.

As it is readily seen from Fig. 4, the DBD-plasma process is more effective when high specific input energy is applied. At $18 \, J \, L^{-1}$, the degradation rate, by DBD-plasma, is around 37%, thus $63.29 \, mg \, m^{-3}$ of Butyraldehyde is measured at the inlet of the fixed-bed. Interestingly, only 17% degradation is achieved with $9 \, J \, L^{-1}$ of SIE.

The concentration of Ozone and BUTY at the outlet of the DBD, fixed-bed reactor and the ratio between Ozone and Butyraldehyde have been determined during the oxidation phase and reported in Fig. 5. An increase of specific input energy from 9 to $18\,\mathrm{J\,L^{-1}}$ enhanced the Ozone decomposition; at $18\,\mathrm{J\,L^{-1}}$, the ratio was about ten times higher than with SIE of $9\,\mathrm{J\,L^{-1}}$. The value of the ratio (O₃/BUTY) at $9\,\mathrm{J\,L^{-1}}$ is 0.46 and it reaches 4.7 for $18\,\mathrm{J\,L^{-1}}$. The difference in the rate of Ozone

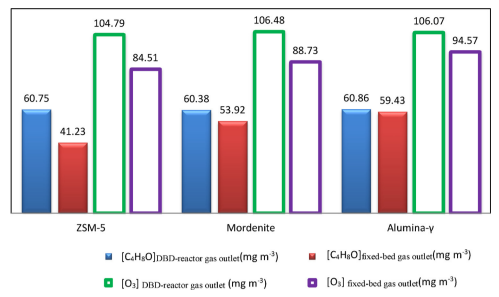


Fig. 3. Ozone and Butyraldehyde concentration, at DBD-reactor outlet, under DBD-plasma/fixed-bed system with three differents catalysts $[Q=1 \text{ m}^3 \text{ h}^{-1}, SIE_{plasma}=18 \text{ J L}^{-1}, [C_4H_8O]_0=100 \text{ mg m}^{-3}, \text{ catalyst}=20 \text{ g}, T=25 ^{\circ}C, RH=45 ^{\circ}M].$

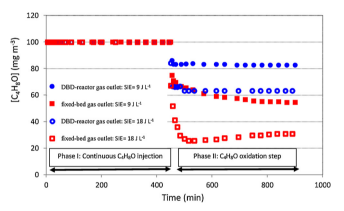


Fig. 4. Variation of C_4H_8O outlet concentration as function of time with differents specific energy under DBD-plasma/fixed-bed system in the case of ZSM-5(Phase I: inlet C_4H_8O injection; Phase II: C_4H_8O oxidation step) $[Q=1 \text{ m}^3 \text{ h}^{-1}, \text{ SIE}_{\text{plasma}}=18 \text{ JL}^{-1}, [ZSM-5]=20 \text{ g}, T=25 ^{\circ}C, HR=5 \%].$

decomposition between the two ratios suggests that all active sites of ZSM-5 are not fully occupied by Ozone at 9 J L⁻¹.

3.2.2. Effect of C_4H_8O concentration

At fixed SIE of 18 JL⁻¹, the effect of BUTY concentration on degradation rate and ozone consumption at the fixed bed reactor has been investigated. The BUTY concentrations are: (i) $100 \, \text{mg m}^{-3}$ and (ii) $200 \, \text{mg m}^{-3}$ with a continuous flowrate of $1 \, \text{m}^3 \, \text{h}^{-1}$ and RH = 5%. The inlet concentration of BUTY significantly influences the ratio of O_3/C_4H_8O). Experimental data for Ozone and Butyraldehyde consumption during C_4H_8O degradation when applying DBD-plasma and post-plasma (fixed-bed) treatments are summarized in Fig. 6. Regarding the influence of inlet concentration, a result shows that the ratio (O_3/C_4H_8O) was significantly affected by increasing C_4H_8O concentration.

For 100 mg m^{-3} of Butyraldehyde at the inlet of (DBD-plasma/fixed-bed) system, the ratio (O₃/BUTY) reaches 4.7 whereas for high concentration ([C₄H₈O] = 200 mg m^{-3}); the ratio is 2.1.

The removal efficiency of C_4H_8O significantly decreased from 51.07%, at $[C_4H_8O] = 100 \, mg \, m^{-3}$, to 16.11% for $[C_4H_8O] = 200 \, mg \, m^{-3}$ (Fig.6). The abatement of Butyraldehyde mediated by the DBD-plasma process as well as on fixed-bed reactor is higher at a lower initial concentration. It appears that for C_4H_8O inlet concentration of 200 mg m⁻³, the air flow is fairly saturated with

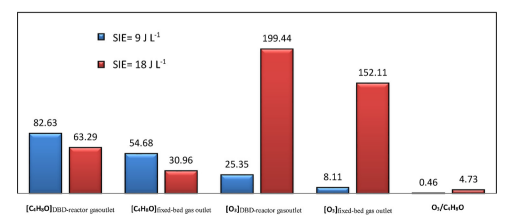


Fig. 5. C_4H_8O and Ozone concentrations (mg m⁻³) in outlet DBD-reactor and outlet fixed-bed for two different specific energy [Q = 1 m³ h⁻¹, [ZSM-5] = 20 g, T = 25 °C, HR = 5 %].

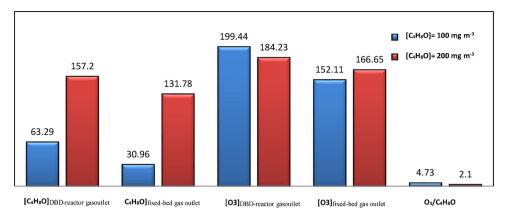


Fig. 6. C_4H_8O and Ozone concentrations (mg m⁻³) in DBD-reactor outlet and fixed-bed outlet for two differents inlet concentrations [Q = 1 m³ h⁻¹, SIE_{plasma} = 18 JL⁻¹, [ZSM-5] = 20 g, T = 25 °C, HR = 5 %].

Table 3 Experimental conditions selected for C_4H_8O oxidation under (DBD-plasma/fixed-bed) system.

Parameter	Q_{air}	$[C_4H_8O]_0$	SIE _{plasma}	W _{ZSM-5}
Value	1 m ³ h ⁻¹	100 mg m ⁻³	18 J L ⁻¹	20 g

Butyraldehyde, which limits the access of Ozone molecules to the micropores of ZSM-5, thus preventing its decomposition. The increase in the amount of Butyraldehyde causes a significant decrease of Ozone and Butyraldehyde. Thus, Ozone cannot be adsorbed on active sites and prevents the reactive sites to intervene in catalytic oxidation. Ozone contributes to oxidize Butyraldehyde already adsorbed on the zeolite and additionally, a part of the ozone is consumed to generate radicals on the free sites. Ozone decomposition occurs at the Lewis sites of zeolite, to produce atomic oxygen, peroxide ions and hydroxyl-radicals according to Eqs. (5)–(7) [22], to oxidize the adsorbed Butyraldehyde.

$$O_3 + Z - Al \rightarrow Z - Al...O + O_2 \tag{5}$$

$$Z-Al...O+O_3 \rightarrow Z-Al...O_2+O_2$$
 (6)

$$Z - Al...O_2 \rightarrow Z - Al + O_2 \tag{7}$$

3.2.3. Effect of humidity

The humidity is inevitably present in the environment, thus the study of the effect of humidity on BUTY removal and Ozone decomposition were monitored in a humid environment. Three experiments, with ZSM-5 catalyst, have been carried out with relative humidity (RH) of 5, 45% and 90%. The optimized experimental conditions used during C_4H_8O degradation are summarized in Table 3.

Experimental results of Ozone and Butyraldehyde consumption rates for C_4H_8O degradation by DBD-plasma and post-plasma treatment under different humidity levels are presented in Fig. 6. The results show that the variation of RH causes a significant variation in terms of Butyraldehyde oxidation on DBD-plasma reactor. Indeed, a good removal efficiency is reached in the case of medium humidity of RH = 45%. However, it is observed that the increase in moisture significantly decreased the Ozone production, in agreement with references [10,11,16,20]. Fig. 7 illustrates the decrease of C_4H_8O degradation efficiency during the catalytic oxidation processes.

The influence of relative humidity on Butyraldehyde degradation under catalytic oxidation was also monitored. By increasing the relative humidity in the air stream, a drop in consumption rates of the residual pollutants, namely O_3 and C_4H_8O , is observed. This was due to the effect of water molecules and Ozone adsorbed on the active sites of ZSM-5. Ozone will undergo a lower decomposition, which lead to a decrease in the performance of Butyraldehyde degradation.

3.2.4. Effect of the catalyst dosage

The influence of weight of the catalyst ZSM-5 on BUTY and Ozone removal is reported in this section. Fig. 8 shows the removal efficiency of BUTY under (DBD-plasma/fixed-bed) system for two catalytic bed presenting different heights. It was observed that the increase in catalyst bed increased the BUTY removal efficiency. It can be seen that at 15 cm, i.e. 60 g of ZSM-5, Ozone concentration measured at the outlet of (DBD-plasma/fixed-bed) system is practically zero ([O₃] = 0.51 mg m $^{-3}$). In addition, BUTY consumption rate consumption rate was higher compared to catalytic beds 5 cm high. A higher catalyst height provides a reaction medium enhancing the Ozone decomposition

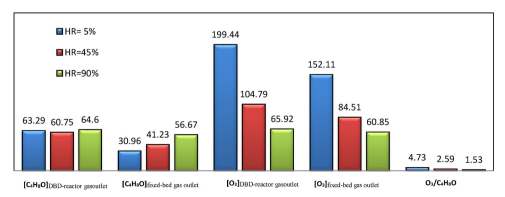


Fig. 7. C_4H_8O and Ozone concentrations (mg m⁻³) in the outlet of DBD-reactor and outlet of fixed-bed with differents humidity (Q = 1 m³ h⁻¹, SIE_{plasma} = 18 JL⁻¹, [ZSM-5] = 20 g, T = 25 °C, HR = 5 %).

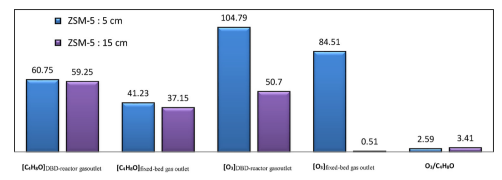


Fig. 8. C_4H_8O and Ozone concentrations (mg m⁻³) in the outlet of DBD-reactor and outlet of fixed-bed for two differents heights [Q = 1 m⁻³ h⁻¹, SIE_{plasma} = 18 JL⁻¹, [ZSM-5] = 20 g, T = 25 °C, HR = 5 %].

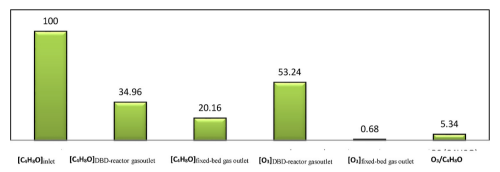


Fig. 9. Variation of C_4H_8O concentration (mg m⁻³) and Ozone formation (mg m⁻³) under UV-lamp/DBD-plasma/fixed-bed system for ZSM-5 [Q = 1 m³ h⁻¹, SIE_{plasma} = 18 JL⁻¹, [ZSM-5] = 20 g, T = 25 °C, HR = 5 %].

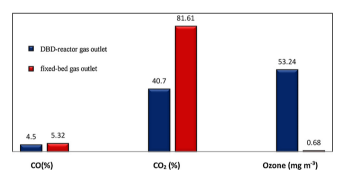


Fig. 10. CO_x selectivity (%) and ozone concentration in the outlet of DBD-reactor and outlet of fixed-bed [Q = $1 m^3 h^{-1}$, [C₄H₈O] = 100 mg m^{-3} , [ZSM-5] = 20 g, SIE_{plasma} = 18 JL^{-1} , T = $25 \, ^{\circ}$ C, HR = $45 \, ^{\circ}$].

Table 4Surface atomic percentage of elements as determined by XPS.

	C1s	N1s	O1s	Al2p	Si2p
Before	9.69	0.45	61.37	22.89	5.61
After	15.04	0.1	58.82	21.89	4.14

 $3.3. \ Effect of combined \ Plasma/Photocatalysis \ system \ and \ Catalytic \\ Ozonation \ on \ Ozone \ removal \ and \ Butyraldehyde \ oxidation$

Process performance (UV-lamp/DBD-plasma/fixed-bed) was monitored during the photocatalysis (TiO₂/UV-lamp) and DBD-plasma process for Butyraldehyde degradation. For this purpose, the UV lamp within DBD-plasma reactor was applied during the 8 hour runs. Coupling plasma technologies with photocatalysis for Butyraldehyde degradation was reported by our laboratory [24,16]. The association of

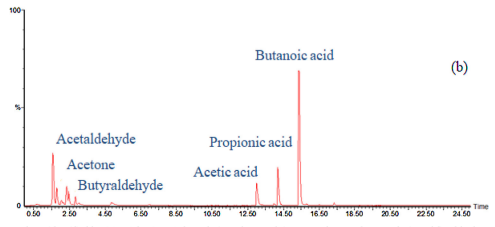


Fig. 11. Intermediate by-products identified by GC-MS for C_4H_8O degradation when applying DBD-plasma/photocatalysis and fixed-bed reactor: (a) DBD-reactor gas outlet: DBD-plasma/photocatalysis, (b) fixed-bed gas outlet.

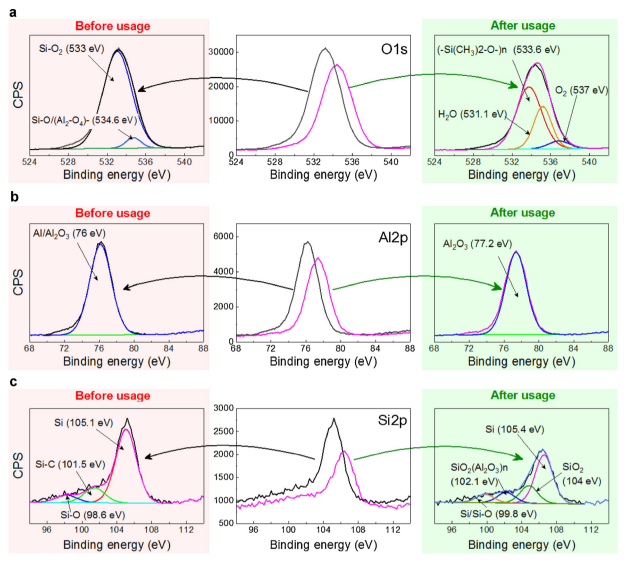


Fig. 12. a) XPS deconvolution of O1s in the ZSM-5 catalytic beds before and after use in BUTY removal in the DBD-plasma/photocatalysis/catalytic ozonation system. b) XPS deconvolution of Al2p in the ZSM-5 catalytic beds before and after use in BUTY removal in the DBD-plasma/photocatalysis/catalytic ozonation system. c) XPS deconvolution of Si2p in the ZSM-5 catalytic beds before and after use in BUTY removal in the DBD-plasma/photocatalysis/catalytic ozonation system.

both technologies in the same process led to synergistic effects improving the efficiency of VOCs removal [23,25-30]. In order to approach the conditions of the industrial application of this process. Processes reported the coupling of DBD-plasma with photocatalysis (TiO₂/UV-lamp) under humid conditions (HR = 45%) applying continuous flow of 1 m³ h⁻¹ and 100 mg m⁻³ of Butyraldehyde. The catalytic bed downstream of (UV-lamp/DBD-plasma) process was loaded with 60 g of ZSM-5 already saturated with Butyraldehyde. C₄H₈O adsorption step was realized on batch reactor (Pyrex 1L) and after reaching the adsorption equilibrium. The catalysts were then placed in the fixed-bed reactor. The results (Fig. 9) show that the amount of Ozone at the outlet of fixed-bed is low and almost zero that is total consumption of ozone was observed on of ZSM-5. In addition, a more effective degradation is observed for Butyraldehyde. The catalytic bed allows complete decomposition of Ozone and good degradation of Butyraldehyde. Ozone concentration at the outlet of (UV-lamp/DBDplasma/fixed-bed) was around 0.68 mg m⁻³ and the rate of Butyraldehyde removal efficiency reached 79.8%.

3.4. Pollutantmineralization (CO and CO_2 selectivity's) and by-products identification

The evolution of CO and CO₂ selectivity and Ozone concentration at the outlet of DBD-plasma/photocatalytic reactor and fixed-bed are presented in Fig. 10. An increase in mineralization rate and a total consumption of Ozone is observed (Fig. 10) during the catalytic oxidation of Butyraldehyde. This observation shows that humidity enhances the decomposition of Ozone on ZSM-5 into hydroxyl radicals and consequently favors mineralization [31–34]. Indeed, CO₂ reached the highest selectivity at the outlet of fixed-bed reactor with a rate of 81.61% against 40.7% for DBD-plasma/photocatalytic reactor and a higher performance in Ozone removal on ZSM-5 [29,35]. Brodu et al., obtained similar results for the toluene degradation [22,28] under catalytic ozonation. The CO, selectivity measured either at the outlet of DBD-plasma/photocatalytic reactor or at the outlet of fixed-bed reactor remained constant around 5% (see Fig. 10).

The by-products generated under DBD-plasma/photocatalytic reactor (DBD-plasma/UV-lamp) and fixed-bed reactor were monitored by Gas Chromatography coupled to Mass Spectrometry (GC-MS). All samples are concentrated on a Carbotrap (25 ml) and then removed by thermal desorption unit coupled with GC-MS. The chromatograms are

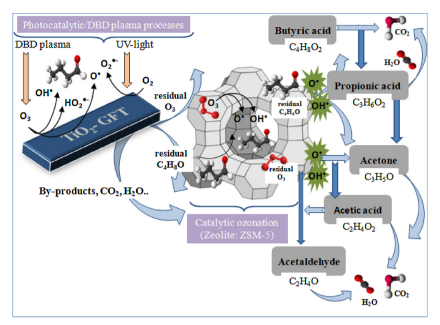


Fig. 13. Suggested mechanism for Butyraldehyde removal under simultaneous DBD-plasma/photocatalysis/catalytic ozonation.

shown in Fig. 11. The intermediate by-products detected other than CO_2 were: acetone (C_3H_5O) , acetaldehyde (C_2H_4O) , acetic acid $(C_2H_4O_2)$, propionic acid $(C_3H_6O_2)$ and butanoic acid $(C_4H_8O_2)$.

The surface atomic concentrations of elements before and after the Ozone and residual VOC decomposition were investigated by XPS. Table 4 shows the zeolite surface atomic concentration percentage after the eff; ect of the combined catalytic processes [25]. During the catalytic ozonation, the carboxylic acid intermediates were partly mineralized to $\rm CO_2$ and $\rm H_2O$ and partly increased the C1s concentration on the catalysts surface as seen in Table 4 from 9.69 to 15.04% during Butyraldehyde degradation in the presence of Ozone [16,20,36–40].

Fig. 12 shows the oxidative states of the elements reported in Table 2 before and after the pollutant removal under light. Before usage of the ZSM-5 beds, the O1s showed peaks at 534.6 and 533 eV reflecting O-Si and O-Al bindings. After usage, ZSM-5 shows the presence of H₂O [41] and O₂ [42], which can be attributed to: (i) the attributed humidity during the pollutant removal, (ii) the removal/mineralization of the pollutants generating CO₂ and H₂O, and (iii) the contact with air after the reaction when recovering the ZSM-5 beds. This later factor did not significantly affect the non-used beds, which favors the first two arguments. Surprisingly, the signal related to Al-O was not detected at this stage. Before use, ZSM-5 presented partially oxidized Al2p. However, after exposure to the photocatalytic oxidation and the generated O₃, the Al2p shows peak at 77.2 reflecting totally oxidized Al, mainly Al₂O₃. When exposed to oxygen, the Al metal surface oxidizes reaching saturation, while the subsurface may stay showing metallic Al, with a characteristic Al2p signal at ~76 eV. The saturated interface reacts further with oxygen, and eventually the bulk metal is transformed into oxide at room temperature. It is concluded that molecular oxygen does not dissociate on the oxide while atomic oxygen is still capable of adsorbing and diffusing into the bulk as recently reported by Cakir et al. [43]. The same authors showed that atomic oxygen adsorption broadens the O2p valence band. All photoemission peaks underwent a rigid shift to low binding energies, which is consistent with the formation of O₂ surface atoms [43]. The deconvolution of Si2p spectra showed the presence of SiO₂ (Al₂O₃)n [44] and Si-C [45].

The peaks shifts observed in Fig. 12 (a, b and c) after the ZSM-5 usage in the catalytic removal of BUTY can be due to the oxidative states of the elements under light. The observed changes in the peaks intensities and shifts in the binding energies (BE) of O1s, Al2p and Si2p provide the evidence for *redox* reactions taking place during the

pollutant degradation time. This is related to the relative sensitivity factors of each element and the observed intensities of all elements on the catalyst surface. It is worth to mention that the peaks intensities depend on the X-ray emission, charge neutralization, signal acquisition resolution and the samples themselves [16,39].

The mechanism for C₄H₈O removal when applying simultaneously DBD-plasma/photocatalysis/catalytic ozonation processes is suggested in Fig. 13. C₄H₈O and O₃ suggested to be adsorbed on the zeolites surface promoting a surface reaction where O3 is decomposed into surface active oxygen species/radicals reacting with Butyraldehyde [38,39]. In our previous studies [16,25] we reported on the degradation of air pollutants by coupling DBD-plasma and photocatalysis without catalytic ozonation using zeolites. These previous studies showed that the degradation of Butyraldehyde is the result of its oxidation by the photogenerated reactive oxygen species (ROS) created by plasma/ photocatalysis (atomic oxygen, peroxide ions and hydroxyl radicals). The major by products were acetic acid (C₂H₄O₂), propionic acid $(C_3H_6O_2)$ and butanoic acid $(C_4H_8O_2)$. In this study, when we added the zeolite fixed-bed reactor at the outlet of DBD-plasma /photocatalysis reactor, the outcomes show that the amount of residual Butyraldehyde decreased at the outlet of the fixed-bed reactor and the amount of the same by-products increased (Fig.11). These results show that a similar mechanism occurs when adding the zeolite fixed-bed reactor i.e. the generation of more ROS due to the contact of the Ozone (coming from plasma/photocatalysis reactor) with the active surface sites of the zeolites and the oxidation of Butyraldehyde in the fixed-bed with the ROS.

4. Conclusion

Butyraldehyde degradation in a DBD-photocatalytic reactor by catalytic oxidation (on fixed-bed reactor) on ZSM-5, Mordenite and γ -alumina was reported. In this study, we show that by adding zeolite fixed-bed reactor to the plasma/photocatalysis reactor, we were able to: (i) achieve higher degradation rate of the residual Butyraldehyde in fixed-bed via the residual Ozone generated from plasma system, and (ii) reduce the amount of ozone generated in the plasma/photocatalysis reactor. The values obtained for SIE_{plasma}, [C_4H_8O]_{inlet}, RH (%) and ZSM-5 height show that ZSM-5 intervened in the oxidation. This was possible due to its porous structure oxidizing the Butyraldehyde adsorbed via the Ozone/reactive species. Under humid conditions, the

strong affinity of ZSM-5 for water molecules lead to a lower catalytic yield in the photocatalysis/fixed-bed reactor. A total reduction of Ozone concentration was attained: $\sim 0,68~\text{mg m}^{-3}$ against 53,24 mg m $^{-3}$ under plasma/photocatalysis process without catalytic ozonation. The system plasma/photocatalysis/ZSM-5 allowed complete elimination of Butyraldehyde with a mineralization rate of 81.61%. This opens doors for possible industrial applications of this system in automotive cabinets, hospitals' indoor odors, and domestic chimneys.

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